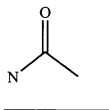
## IN THE CLAIMS

1. (Currently amended) A bisubstrate inhibitor of insulin receptor kinase, comprising:

a nucleotide or nucleotide analog moiety comprising a triphosphate
consisting of γ-S-ATP; and

a peptide moiety which is a substrate for said insulin receptor kinase and which comprises a tyrosine residue or a 2-amino-3-(4-amino-phenyl)-propionic acid residue;

wherein said moieties are linked by a tether, wherein said tether is linked to the tyrosine residue via its phenolic oxygen or to the 2-amino-3-(4-amino-phenyl)-propionic acid residue via its aniline nitrogen and wherein said tether is linked to the nucleotide or nucleotide analog moiety via the gamma phosphate of said  $\gamma$ -S-ATP the triphosphate, wherein the tether is



greater than or equal to 4.9 Å measured from a gamma phosphorus of the nucleotide or nucleotide analog moiety to a proton donor of the tether formed by the phenolic oxygen or the aniline nitrogen.

- 2. (Cancelled)
- 3. (Cancelled)
- 4. (Cancelled)
- 5. (Currently amended) The bisubstrate inhibitor of claim 1 wherein the peptide moiety has at least 4 contiguous amino acid residues selected from the sequence Lys Lys Lys Leu Pro Ala Thr Gly Asp Tyr Met Asn Met Ser Pro Val Gly Asp (SEQ ID NO:1), wherein the Tyr residue is modified so that a nitrogen atom replaces a hydroxyl to form the 2-amino-3-(4-amino-phenyl)-propionic acid residue.
- 6. (Currently amended) The bisubstrate inhibitor of claim 1 wherein the peptide moiety has at least 5 contiguous amino acid residues selected from the sequence Lys Lys

- Lys Leu Pro Ala Thr Gly Asp Tyr Met Asn Met Ser Pro Val Gly Asp (SEQ ID NO:1), wherein the Tyr residue is modified so that a nitrogen atom replaces a hydroxyl to form the 2-amino-3-(4-amino-phenyl)-propionic acid residue.
- 7. (Currently amended) The bisubstrate inhibitor of claim 1 wherein the peptide moiety comprises the sequence Lys Lys Lys Leu Pro Ala Thr Gly Asp Tyr Met Asn Met Ser Pro Val Gly Asp (SEQ ID NO:1), wherein the Tyr residue is modified so that a nitrogen atom replaces a hydroxyl to form the 2-amino-3-(4-amino-phenyl)-propionic acid residue.
- 8. (Cancelled)
- 9. (Cancelled)
- 10. (Original) The bisubstrate inhibitor of claim 1 wherein the peptide moiety comprises a membrane translocating sequence (MTS).
- 11. (Original) The bisubstrate inhibitor of claim 10 wherein the MTS is at the N-terminus of the peptide moiety.
- 12. (Original) The bisubstrate inhibitor of claim 10 wherein the MTS is at the C-terminus of the peptide moiety.
- 13. (Original) The bisubstrate inhibitor of claim 1 wherein the peptide moiety comprises an HIV TAT sequence.
- 14. (Cancelled)
- 15. (Currently amended) A bisubstrate inhibitor of insulin receptor kinase, emprising:

  a nucleotide or nucleotide analog moiety;

and a peptide moiety which is a substrate for said insulin receptor kinase; wherein said moieties are linked by a tether that comprises a proton donor; wherein the tether is greater than or equal to 4.9 Å measured from a gamma phosphorus of the nucleotide or nucleotide analog moiety to the proton donor; wherein the bisubstrate inhibitor of insulin receptor kinase is Compound 2.

- 16-57. (Cancelled)
- 58. (Original) The bisubstrate inhibitor of claim 1 which is bound to insulin receptor kinase.
- 59. (Cancelled)
- 60. (Currently amended) A bisubstrate inhibitor of a protein kinase comprising:

a nucleotide or nucleotide analog moiety comprising a triphosphate consisting of  $\gamma$ -S-ATP; and

a peptide moiety which is a substrate for said protein kinase and which comprises a-tyrosine residue, a 2-amino-3-(4-amino-phenyl)-propionic acid residue or ; a serine residue, a 2,3-diamino-propionic acid residue, a threonine residue, or a 2,3-diamino-butyrie acid residue;

wherein said moieties are linked by a tether, wherein said tether is linked to the tyrosine residue via its phenolie exygen, to the 2-amino-3-(4-amino-phenyl)-propionic acid residue via its aniline nitrogen, to the serine residue via its hydroxyl exygen, or to the 2,3,diamino-propionic acid residue via its 3-amino nitrogen, to the threonine residue via its hydroxyl exygen, or to the 2,3-diamino-butyrie acid via its 3-amino nitrogen, and wherein said tether is linked to the nucleotide or nucleotide analog moiety via the gamma phosphate of said γ-S-ATP the triphosphate, wherein the tether is



greater than or equal to 4.9 Å measured from a gamma phosphorus of the nucleotide or nucleotide analog to a proton donor of the tether formed by the phenolic oxygen, the aniline nitrogen, the hydroxyl oxygen, or the 3-amino nitrogen.

- 61. (Cancelled)
- 62. (Cancelled)
- 63. (Currently amended) The bisubstrate inhibitor of claim 60 wherein the protein kinase is a tyrosine protein kinase and the peptide comprises a tyrosine residue.

  wherein the tyrosine residue is modified so that a nitrogen atom replaces a hydroxyl to form the 2-amino-3-(4-amino-phenyl)-propionic acid residue.
- 64. (Cancelled)
- 65. (Cancelled)

- 66. (Cancelled) The bisubstrate inhibitor of claim 63 wherein a nitrogen atom replaces a hydroxyl oxygen on the tyrosine.
- 67. (Original) The bisubstrate inhibitor of claim 60 which is bound to the protein kinase.
- 68. (Cancelled)
- 69. (Currently amended) The bisubstrate inhibitor of claim 60 wherein the peptide moiety comprises at least 4 contiguous amino acids of a natural substrate of said protein kinase wherein the peptide moiety comprises either a tyrosine residue that is modified so that a nitrogen atom replaces a hydroxyl to form the 2-amino-3-(4-amino-phenyl)-propionic acid residue or a serine residue that is modified so that a nitrogen atom replaces a hydroxyl to form a 2,3-diamino-propionic acid residue.
- 70. (Currently amended) The bisubstrate inhibitor of claim 60 wherein the peptide moiety comprises at least 5 contiguous amino acids of a natural substrate of said protein kinase, wherein the peptide moiety comprises either a tyrosine residue that is modified so that a nitrogen atom replaces a hydroxyl to form the 2-amino-3-(4-amino-phenyl)-propionic acid residue or a serine residue that is modified so that a nitrogen atom replaces a hydroxyl to form a 2,3-diamino-propionic acid residue.
- 71. (Currently amended) The bisubstrate inhibitor of claim 60 wherein the peptide moiety comprises at least 6 contiguous amino acids of a natural substrate of said protein kinase, wherein the peptide moiety comprises either a tyrosine residue that is modified so that a nitrogen atom replaces a hydroxyl to form the 2-amino-3-(4-amino-phenyl)-propionic acid residue or a serine residue that is modified so that a nitrogen atom replaces a hydroxyl to form a 2,3-diamino-propionic acid residue.
- 72. (Cancelled)
- 73. (Cancelled)
- 74. (Currently amended) The bisubstrate inhibitor of claim 60 wherein the peptide moiety is a natural substrate of said protein kinase, wherein the peptide moiety comprises either a tyrosine residue that is modified so that a nitrogen atom replaces a hydroxyl to form the 2-amino-3-(4-amino-phenyl)-propionic acid residue or a serine residue that is modified so that a nitrogen atom replaces a hydroxyl to form a 2,3-diamino-propionic acid residue.
- 75. (Cancelled)

76. (Cancelled)

77. (New) A method of making a candidate bisubstrate inhibitor of insulin receptor kinase, comprising:

synthesizing a peptide comprising a nitrophenylalanine residue; reducing the nitro group on said nitrophenylalanine residue to form an amine group;

bromoaceteylating the amine group to form a bromide group; displacing the bromide group by a phosphorothicate of ATPyS

78. (New) A method of making a candidate bisubstrate inhibitor of a protein kinase comprising:

synthesizing a peptide comprising a residue selected from the group consisting of a nitrophenylalanine residue and a diamino propionic acid residue; reducing the nitro group on said nitrophenylalanine residue to form an amine group, if the peptide comprises a nitrophenylalanine residue; bromoacetylating the amine group on said residue to form a bromide group; displacing the bromide group by a phosphorothioate of ATPγS

79. (New) The method of claim 77 wherein the peptide is selected from the group consisting of:

KKKLPATGD-nitrophenylalanine-MNMSPVGD,

TGD-nitrophenylanaine,

GD-nitrophenylanaine-M,

D-nitrophenylanaine-MN,

nitrophenylanaine-MNM,

ATGD-nitrophenylanaine,

TGD-nitrophenylanaine-M,

GD-nitrophenylanaine-MN,

D-nitrophenylanaine-MNM,

nitrophenylanaine-MNMS,

PATGD-nitrophenylanaine,

ATGD-nitrophenylanaine-M,

TGD-nitrophenylanaine-MN,

GD-nitrophenylanaine-MNMS, D-nitrophenylanaine-MNMSP, LPATGD-nitrophenylanaine-M, ATGD-nitrophenylanaine-MN, TGD-nitrophenylanaine-MNMS, GD-nitrophenylanaine-MNMS, D-nitrophenylanaine-MNMSP, and nitrophenylanaine-MNMSPV.

- 80. (New) The method of claim 77 further comprising the step of testing the candidate bisubstrate inhibitor in kinase assays to determine potency of inhibition.
- 81. (New) The method of claim 77 further comprising the step of testing the candidate bisubstrate inhibitor in kinase assays to determine specificity of inhibition.
- 82. (New) The method of claim 78 wherein the peptide is selected from the group consisting of: known phosphorylation sites of kinase enzymes, wherein a nitrophenylalanine is substituted for a tyrosine residue or a diaminopropionic acid residue is substituted for a serine residue.
- 83. (New) The method of claim 78 wherein the peptide is selected from the group consisting of:

KKKLPATGD-nitrophenylalanine-MNMSPVGD,

TGD-nitrophenylanaine,

GD-nitrophenylanaine-M,

D-nitrophenylanaine-MN,

nitrophenylanaine-MNM,

ATGD-nitrophenylanaine,

TGD-nitrophenylanaine-M,

GD-nitrophenylanaine-MN,

D-nitrophenylanaine-MNM, nitrophenylanaine-MNMS, PATGD-nitrophenylanaine, ATGD-nitrophenylanaine-M, TGD-nitrophenylanaine-MN, GD-nitrophenylanaine-MNM, D-nitrophenylanaine-MNMS, nitrophenylanaine-MNMSP, LPATGD-nitrophenylanaine, PATGD-nitrophenylanaine-M, ATGD-nitrophenylanaine-MN, TGD-nitrophenylanaine-MNM, GD-nitrophenylanaine-MNMS, D-nitrophenylanaine-MNMSP, nitrophenylanaine-MNMSPV, LRRA-diaminopropionic acid-LG, RRA-diaminopropionic acid, RA-diaminopropionic acid-L, A-diaminopropionic acid-LG, LRRA-diaminopropionic acid, RRA-diaminopropionic acid-L, LRRA-diaminopropionic acid-L, and RRA-diaminopropionic acid-LG.

- 84. (New) The method of claim 78 further comprising the step of testing the candidate bisubstrate inhibitor in kinase assays to determine potency of inhibition.
- 85. (New) The method of claim 78 further comprising the step of testing the candidate bisubstrate inhibitor in kinase assays to determine specificity of inhibition.

- 86. (New) The method of claim 77 wherein the peptide is selected from the group consisting of: known phosphorylation sites of insulin kinase enzymes, wherein a nitrophenylalanine is substituted for a tyrosine residue.
- 87. (New) An inhibitor of insulin receptor kinase, comprising: a nucleotide analog moiety consisting of γ-S-ATP; and a peptide moiety which comprises a 2-amino-3-(4-amino-phenyl)propionic acid residue;

wherein said moieties are linked by a tether, wherein said tether is linked to the 2-amino-3-(4-amino-phenyl)-propionic acid residue via its aniline nitrogen and wherein said tether is linked to the nucleotide analog moiety via the gamma phosphate of said  $\gamma$ -S-ATP , wherein the tether is



88. (New) An inhibitor of a protein kinase comprising:

a nucleotide analog moiety consisting of  $\gamma$ -S-ATP; and a peptide moiety which comprises a 2-amino-3-(4-amino-phenyl)-propionic acid residue or a 2,3-diamino-propionic acid residue;

wherein said moieties are linked by a tether, wherein said tether is linked to the 2-amino-3-(4-amino-phenyl)-propionic acid residue via its aniline nitrogen, or to the 2,3,diamino-propionic acid residue via its 3-amino nitrogen, and wherein said tether is linked to the nucleotide analog moiety via the gamma phosphate of said  $\gamma$ -S-ATP, wherein the tether is

